

Virtual reality as an analgesic for acute and chronic pain in adults: a systematic review and meta-analysis

This article was published in the following Dove Press journal:
Journal of Pain Research

Brian Mallari¹
Emily K Spaeth²
Henry Goh³
Benjamin S Boyd⁴

¹Department of Physical Therapy, St. Mary's Medical Center, San Francisco, CA, USA; ²Castro Valley Yoga, Castro Valley, CA, USA; ³Department of Physical Therapy, Kaiser Permanente, Oakland, CA, USA; ⁴Department of Physical Therapy, Samuel Merritt University, Oakland, CA, USA

Background: Previous studies have shown that virtual reality (VR) is effective in reducing acute and chronic pain both in adults and in children. Given the emergence of new VR technology, and the growing body of research surrounding VR and pain management, an updated systematic review is warranted.

Purpose: The purpose of this systematic review is to compare the effectiveness of VR in reducing acute and chronic pain in adults.

Data Sources: A search was conducted in three databases (PubMed, CINAHL, Trip) using standardized search terms.

Study Selection: Twenty experimental and quasi-experimental trials published between January 2007 and December 2018 were included based on prespecified inclusion and exclusion criteria. Pain intensity was the primary outcome.

Data Extraction: We extracted data and appraised the quality of articles using either the PEDro or Modified Downs and Black risk of bias tools.

Data Synthesis: The majority of studies supported the use of VR to reduce acute pain both during the procedure and immediately after. Numerous studies found VR reduced chronic pain during VR exposure but there is insufficient evidence to support lasting analgesia. There was considerable variability in patient population, pain condition and dosage of VR exposure.

Limitations: Due to heterogeneity, we were unable to perform meta-analyses for all study populations and pain conditions.

Conclusions: VR is an effective treatment for reducing acute pain. There is some research that suggests VR can reduce chronic pain during the intervention; however, more evidence is needed to conclude that VR is effective for lasting reductions in chronic pain.

Keywords: virtual reality, analgesia, acute, chronic, pain management, adult

Plain language summary

Virtual reality (VR) could potentially help reduce pain in adults undergoing medical procedures or suffering with chronic pain. We performed a systematic review of the available research to see if VR is helpful at reducing pain. We found twenty research studies of various quality published from January 2007 to December 2018. Ten studies examined VR for acute pain conditions, including pain related to burns or medical procedures. Ten studies examined chronic pain, including musculoskeletal, neuropathic, mixed musculoskeletal and neuropathic or unspecified chronic pain conditions. Most of the studies found that VR helped to decrease acute pain during and immediately after various medical procedures. Some studies found VR reduced chronic pain when people were using the VR device and immediately

Correspondence: Benjamin S Boyd
Department of Physical Therapy, Samuel Merritt University, 450 30th Street, Suite 3728, Oakland, CA 94609, USA
Tel +1 510 879 9200 x7376
Fax +1 510 457 2112
Email bboyd1@samuelmerritt.edu

after but the decrease in pain did not last with the protocols tested to date. However, more research is needed to determine if VR can be effective for lasting reductions in people with chronic pain.

Introduction

Pain management is a growing health-care concern in the US. An estimated 100 million adults in the US suffer from chronic pain, and approximately \$17.8 billion is spent prescribing pain medication annually.¹ Historically, opioid medications have been used for pain management, but there are negative side effects associated with opioid use, including a potential delay in recovery and increased risk of permanent disability.² In addition, opioid use has been associated with increased hospital admissions, increased health-care costs, and deaths.² Even with these risks and health-care burdens, the US consumes 80% of the world's opioids.² Alternative pain management strategies are important to consider in an effort to mitigate some of these risks.

A growing body of evidence supports VR as an alternative strategy for management of acute, burn, and experimental pain in adults and children.^{3–11} Additional experiments demonstrated the positive effect of virtual reality (VR) on pain during a variety of medical procedures including chemotherapy^{12–14} and wound care.^{15,16} Other studies also support VR as an adjunct to pain medications.^{17,18} Furthermore, evidence from systematic reviews^{17–21} conclude that VR is effective in the management of acute pain associated with medical procedures, wound debridement, and experimental pain.

Despite emerging evidence, consensus on the effectiveness of VR on chronic pain remains inconclusive. The most recent systematic review on VR both in acute and in chronic pain was performed over 8 years ago.¹⁹ A number of the studies included in that review are less applicable to the clinical setting because they explored the impact of VR on experimental pain, including painful thermal and pressure stimulation. Furthermore, much of the research has focused on children,^{5,7,11,16} which limits generalizability to adult populations. Finally, VR technology can be expensive and nonportable. However, the emergence of more affordable devices such as head mounted displays (HMD) has made VR more feasible for clinical use. Given these concerns, an updated systematic review comparing acute and chronic pain in adults is warranted.

The purpose of this systematic review is to compare the effectiveness of VR in reducing acute and chronic pain in adults compared to standard care without VR or to a sham treatment of nonVR.

Materials and methods

After preliminary searches, we performed our search using the combined terms “Virtual reality AND distraction”, “Virtual reality AND pain”, and “Virtual reality AND analgesia.” It is hypothesized that distraction is a key mechanism for the effectiveness of VR, as the virtual environment consumes enough cognitive capacity to distract the user from their pain.^{3,22} Therefore, the term “distraction” was used to address the user's engagement in their virtual environment and potential mechanism of VR therapy, while the term “analgesia” was used to address pain reduction. Terms such as “immersive” were not included in our search strategy to allow a greater breadth of search as some studies did not make the distinction between immersive and nonimmersive VR. We performed searches using three databases: Turning Research into Practice (TRIP), CINAHL, and MEDLINE (via PubMed). No searches were performed to seek out gray literature. Full details of the search strategy and search filters used are provided in [Appendix A](#).

For the purposes of this review, pain lasting shorter than three months was categorized as acute pain while ongoing, persistent pain lasting three months or longer was categorized as chronic pain.^{23,24} Researchers have defined VR as the interaction between a participant and a simulated three-dimensional, immersive world.¹⁹ Previous research has compared immersive to nonimmersive VR and found immersive VR to be more effective in pain management.^{7,17–20,25–27} Thus, an inclusion criteria for this systematic review was “immersive” VR, defined by the participant being engaged in a simulated world in the form of visual and auditory feedback with the ability to interact with and/or elicit a reaction from their virtual environment. “Nonimmersive” VR and other distraction-based interventions, such as visual illusions, were exclusion criteria for this systematic review. This systematic review was registered in PROSPERO (CRD42018117881).

We screened articles in three stages. First, we screened titles and abstracts of each article and included them for full-text analysis based on the following criteria: written in English, published from January 2007 to December 2018, studied adults (18 years or older), used immersive VR, and used pain intensity as a primary outcome measure. Studies

with a control group were included if the comparison treatment included standard care without VR or a sham treatment of nonVR. Nonexperimental studies were excluded, in addition to studies that used experimental pain, did not incorporate VR, or used VR for purposes outside of pain control. We excluded studies published before 2007 to attain the most current evidence using modern VR technologies.¹⁹

Second, two researchers evaluated the full text of each article to further determine inclusion or exclusion. The references of previously published systematic reviews on VR were also screened for articles that may have been missed in our search strategies. We excluded articles that did not meet the inclusion criteria. Third, if the decision to include or exclude was not clear, we flagged the article for further evaluation. We discussed each flagged article and eligibility was determined based on the overall efforts of the experimenters to (a) create an environment with visual, auditory, and tactile feedback, and (b) provide methods of interaction between the user and virtual environment. Any flagged articles that did not meet these considerations were excluded.

We extracted the following data from included articles: study design, population size and description, acute or chronic pain, outcome measure for pain (numeric rating scale, NRS; visual analog scale, VAS; graphic rating scale, and brief pain inventory), type of pain (musculoskeletal, neuropathic, burn, or procedural), and parameters of VR intervention including definition, description, condition duration, and comparison intervention (as applicable). We also extracted relevant quantitative data required to determine effect size (ES) for studies that had a comparison group. No ES was calculated for single arm studies. We evaluated the homogeneity of included studies and determined the appropriateness of subgrouping and pooling studies within this systematic review.

The primary outcome of interest was pain intensity, subgrouped into pain reported during the intervention or after the intervention for acute compared to chronic pain. For this analysis, data from between-group design studies (randomized controlled trials including cross over designs, and quasi-experimental studies with a comparison group) was utilized to compare the efficacy of VR to reduce pain compared to standard care or a sham VR treatment. Subgroupings were characterized as follows: acute pain during intervention, acute pain after intervention, chronic pain during intervention, and chronic pain after intervention. As a secondary analysis, studies presenting data of

within-group pain intensity reductions were compared to determine expected magnitude of change from VR intervention in patients with acute and chronic pain.

For the meta-analysis, we included studies that used quantitative outcome measures for pain such as the NRS or VAS and that contained enough data to calculate an ES. We then calculated the percentage of total variation across studies that was due to heterogeneity rather than chance (I^2).²⁸ If the pooled data was relatively heterogeneous (50% or more), a random effects model was used, and if it was relatively homogenous (less than 50%) a fixed effects model was used.²⁸

In masked pairs we reviewed each study for risk of bias using the PEDro tool for randomized controlled trials and Modified Downs and Black (MD&B) quality index for nonrandomized studies.^{29,30} We discussed discrepancies, and a third reviewer helped to determine a final score. PEDro scores of 6–10 are considered high quality, 4–5 fair quality, and scores less than or equal to 3 are of poor quality.³¹ Scores of less than 14 on the MD&B are suggested to be of lower quality.³²

Results

A total of 485 articles were identified through database searches (Figure 1). After titles and abstracts were screened, 49 articles were eligible for full text review. After reviewing, a total of 20 articles were included in this systematic review (Table 1). Fourteen of the articles were randomized controlled trials (nine of which were crossover designs), while the other six were quasi-experimental studies (no comparison group). Ten studies examined VR for acute pain conditions, including three in pain related to burns (BP) and seven in medical procedure related pain (MPRP). Ten studies examined chronic pain, including three in musculoskeletal pain conditions (MSKP), four in neuropathic pain conditions (NP), one in a mixture of musculoskeletal and neuropathic pain conditions (MSKP-NP) and two in an unspecified pain condition (UnP). Results of the quality assessments demonstrated that all studies included in this systematic review were of fair to high quality (Table 1).

The VR equipment varied between studies. Eighteen studies used goggles/glasses or another form of HMD that displayed a virtual environment. One study used two 3D projectors (chronic NP), while another used a 2D screen but with dynamic visual cues that responded to changes in treadmill speed (chronic MSKP).

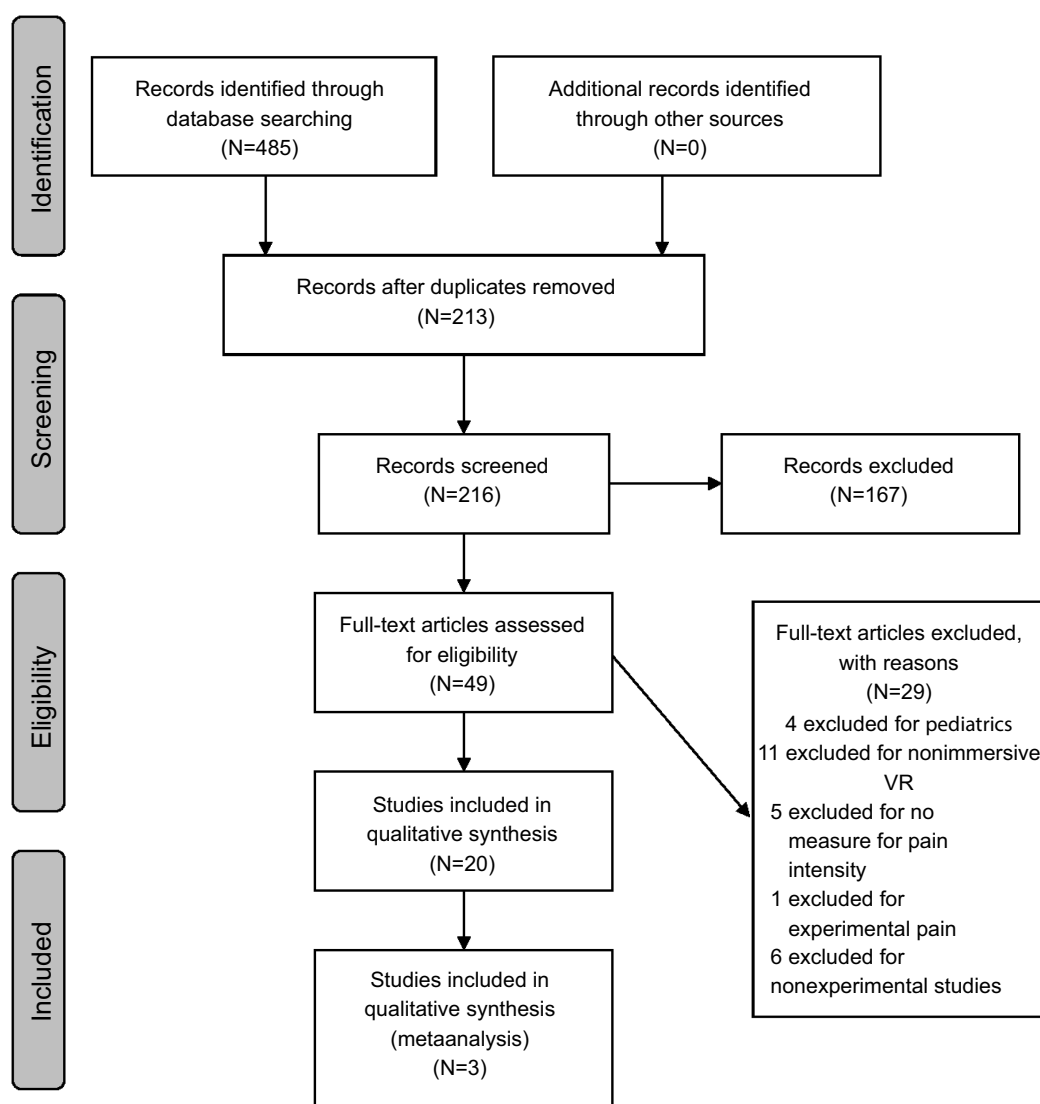


Figure 1 PRISMA flow diagram.
Abbreviation: VR, virtual reality.

Acute pain conditions

Impact of VR on pain experienced during procedure

Three studies examined BP and found that VR in conjunction with medications significantly reduced pain during wound debridement or remobilization of range of motion for limbs affected by burns compared to medications alone.^{9,25,33} Each of these studies used a HMD for VR application and had dosage between 6 and 20 minutes. Pooling of these studies was clinically appropriate due to the homogeneity of the study population and VR application. One study subdivided their sample into two groups based upon severity of worst pain (either $\geq 7/10$ or $< 7/10$).²⁵ These independent groups were pooled together along with data from Carrougher,⁹ for the meta-analysis (Figure 2). However, there was insufficient data to

calculate an ES from Morris et al,³³ thus, this study was not included in the meta-analysis. The pooled ES was 0.66 (95%CI: 0.20, 1.12) (Figure 2). Quality assessment scores for these studies were between 6–7 and 10 on the PEDro scale indicating high quality and low risk of bias.

Four studies evaluated the impact of VR on pain during various medical procedures (MPRP).^{34–37} One study found VR significantly reduced pain during periodontist procedures compared to both a control (standard care) (ES=0.95, 95%CI: 0.64, 1.26) and watching a movie during the procedure (ES=0.44, 95%CI: 0.23, 0.65).³⁴ Another study found VR significantly reduced pain during multiple stages of episiotomy repairs (hymen repair ES=0.74, 95%CI: 0.43, 1.04 and skin repair ES=1.00, 95%CI: 0.65, 1.36) in women after having their first baby compared to repair with no VR.³⁵

Table 1 Individual study characteristics

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Carrougher, 2009 ⁹	RCT, within subjects cross over design	39	Range: 21-57	Inpatients with a mean total body surface area (TBSA) burn of 18% (range, 3–60%)	BP	Acute	PEDro 6/10	Head-position tracked, medical care environment excluding VR helmet with stereophonic sound (Nvis Nvisor).	Patients glided through an icy 3-dimensional canyon with a river and waterfall. They could shoot snowballs (by pressing the spacebar) at snowmen, igloos, and penguins. Contact between the snowball and object or creature elicited a sound, heard only by the patient	10 minutes	Same number of repetitions and same exercises performed in the same plane by the same Physical Therapist without VR. Medications provided for both groups as per standard care.	VR reduced GRS scores for worst pain, pain unpleasantness, and time spent thinking about pain, relative to the no VR condition (27, 31, and 37%, respectively).	Worst pain during treatment	0.54	0.31	0.77	0.004

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Maani, 2011 ²⁵	RCT, within subjects cross over design	6	Range: 20-27	US soldiers burned in combat attacks involving explosive device. Worst pain $\geq$ 7/10.	BP	Acute	PEDro 7/10	Voodoo Envoy laptop. Participants wore a pair of Rockwell Collins SR-80A VR goggles which blocked patient's view of the real world. The goggles were held in place near the patient's eyes by a custom made robotic arm goggle holding system.	Patients "looked" around the virtual environment of an icy canyon with an icy river and heard sound effects (i.e. a splash when a snowball hit the river) mixed with background music. They used a mouse to aim snowballs at various creatures.	6 minutes	Standard care pharmacology without VR.	Worst pain experienced during the treatment procedure for BP was significantly lower during VR compared to the control condition. Time thinking about pain dropped from "most of the time" to "some of the time." Unpleasantness dropped from "mod" to "mild" and worst pain from "mod" to "mild". Increased amount of "fun" with VR during wound care. Significant change in patients with severe pain ($\geq$ 7/10).	Worst pain during treatment	1.51	0.52	2.49	0.043

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
	—	6	Range: 20-27	US soldiers burned in combat attacks involving explosive device. Worst pain < 7/10.	—	—	—	—	—	—	—	No significant changes seen in worst pain reported nor unpleasantness of procedures for patients with mild/mod pain (<7/10). Significant reduction in time thinking about pain and increase in "fun" reported.	Worst pain during treatment	0.43	-0.09	0.96	NS

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/ Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Morris, 2010 ³³	RCT, within subjects cross over design	11	Range: 23-54	Adult burn patients admitted to the Tigerberg Hospital adult burn unit, South Africa. The median Total Body Surface Area (TBSA) was 15% (range 2-55%).	BP	Acute	PEDro 7/10	Low-cost: ASUS F55L Business Laptop and LOGIC PC ATTACK 3 joystick for input and a HMD (eMagin Z800 3D visor)	Patients played "Walt Disney's Chicken Little" PC game with immersive visual feedback and ability to interact with the game via joystick.	20 minutes each (VR plus meds or just meds)	Same number of repetitions and same exercises performed in the same plane by the same Physical Therapist without VR. Medications provided for both groups as per standard care.	No significant difference could be detected between the two sessions ($p = 0.13$; MD = 2.09; 95% CI: -0.67 to 4.85)	Pain during treatment.	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Frey, 2018 ³⁷	RCT, within subjects cross over design	27	Range: 19-38	Eligible patients were otherwise healthy women at ≥32 weeks' gestation giving birth for the first time and in the first stage of labor with an anticipated vaginal delivery. Subjects were recruited from Michigan Medicine's Von Voigtlander Women's Hospital.	MPRP	Acute	PEDro 6/10	Samsung Gear VR HMD powered by a Galaxy S7 phone. User input consisted of head tracking and a hand control	Each patient experienced the same scene of curious manatees from the Ocean Rift (www.oceanrift.com) scuba diving simulation with sounds of manatee calls and breathing underwater. Additional relaxing music was supplied from nighttime sleep by Brain.fm (www.brain.fm). Hand controls simulated taking underwater photos. The participant was observed with VR during unmedicated contractions.	10 minutes	Unmedicated contractions without VR	The numeric rating scale scores of the primary outcome for worst pain intensity were significantly lower in the VR condition (slope estimate −1.5 [95% CI, −0.8 to −2.2] and standardized mean difference −0.8).	Pain during contractions	unable to calculate	unable to calculate	unable to calculate	

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Furman, 2009 ³⁴	Within subjects cross over design (conditions arbitrarily assigned)	38	AVG 45.9 (SD 12.6)	Adults with mild, mod, or severe periodontitis who needed scaling and root planing in all four quadrants	MPRP	Acute	MD&B 22/27	V8 HMD with a Silicon Graphics Octane/MXE workstation with Octane Channel Option	Patients explored a botanical garden in Second Life, a virtual world accessible via the internet, using a handheld mouse. Patients chose their own pathways through the VR environment by controlling the direction of the gaze of the avatar (ex: patients could choose to walk or fly through the garden)	20 minutes	Control (no distraction)	VR condition resulted in significantly lower reported pain level compared to control condition.	Worst pain during treatment	1.80	1.49	2.11	<0.001
	—	—	—	—	—	—	—	—	—	—	Watching a movie (1st 20mins of Cars movie).	VR condition resulted in significantly lower reported pain level compared to watching movie condition.	Worst pain during treatment	0.44	0.23	0.65	0.019

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/ Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Guo, 2015 ³⁸	RCT	98	Experimental group: AVG 30.1 (SD 19.5), Control group: AVG 32.1 (SD 17.4)	Patients with hand injury that needed dressing changes. Ages 18-65. Debridement or suturing within 72 hours of injury.	MPRP	Acute	PEDro 7/10	3D glasses that had a fullbracketing Ruanjiao streamline design to cover the entire eye socket. Headphones were used for sound output; a mouse was used for input.	Patients were immersed in the 3D film "Afanda," which depicts a mysterious dream planet Afanda in which users can reach out to touch a graceful scene.	Patients were asked to watch 3D movies for 5 minutes before the dressing change ended.	Conventional dressing repose with no VR.	The VAS score at the end of the dressing change was significantly lower in the experimental group than in the control group ($t = -30.792$, $p < 0.001$).	Pain immediately after procedure	1.87	1.49	2.25	<0.001

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/ Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
JahanShoorab, 2015 ³⁵	RCT	30	AVG 24.1 (SD 4.1), range: 18-34	Iranian primiparous parturient women having labor at Omolbanin Hospital (ages 18-34).	MPRP	Acute	PEDro 6/10	3D Blu-ray/ DVD player full HD connected to a pair of video glasses (Wrap 920 system, Vizix factory) which include two miniature LCD viewing screens (for the right and left eyes). Two external head-phones were used. Video glasses with audio were worn during episiotomy repair. All repairs were performed by the same expert midwife.	Patients were immersed in the 3D film "IMAX Dolphins and Whales 3D 1080p."	Average of 11.4 minutes (VR) and 13.6 minutes (control)	Standard episiotomy repair without VR.	A significant difference was found between the groups, based group effect ($P=0.038$) and different stages ($P<0.0001$). The pattern of findings, was statistically significant for the pain intensity (group and stages $P=0.044$).	Pain during procedure (Hymen repair)	0.74	0.43	1.04	0.038
	—	—	—	—	—	—	—	—	—	—	—	—	Pain during procedure (Skin repair)	1.00	0.65	1.36	0.038

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
	—	—	—	—	—	—	—	—	—	—	—	—	Pain immediately after procedure	1.37	0.95	1.79	0.038
	—	—	—	—	—	—	—	—	—	—	—	—	Pain 1 hour after procedure	0.62	0.34	0.90	0.038
McSherry, 2018 ⁴⁰	RCT, within subjects cross over design	18	AVG 38.4 (SD 15.5)	Adults patients undergoing painful wound care procedures for deep or partial thickness burns $\geq 5\%$ or complex nonburn wounds, such as necrotizing fasciitis or large decubitus ulcers	MPRP	Acute	PEDro 7/10	NVISINC MX 90 virtual reality goggles and noise-cancelling earphones.	Patients engaged in "Snow World," a virtual reality environment where the participant is tasked with throwing snowballs at objects by clicking a computer mouse button. Music and sound effects from "Snow World" were used.	Minutes for dressing change with VR was 29.9 ± 12.9 . Minutes for dressing change without VR was 30.7 ± 15.1 .	Dressing procedure without VR	No significant difference was found in pain reduction scores between the IVR (Immersive Virtual Reality) treatment group and the No IVR treatment group ($P>0.05$). The IVR treatment lead to reductions in pain of 1.2 ± 2.9 , while the No IVR treatment lead to reductions in pain of 0.3 ± 1.7 .	Pain immediately after procedure	0.41	0.09	0.74	<0.05

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Mosso-Vasquez, 2014 ³⁹	Quasi-experimental study (no comparison group)	67	Not reported	Patients within 24 hours post cardiac surgery (valve replacement, revascularization, stent insertion, tricuspid plasty, communication repair, tricuspid resection).	MPRP	Acute	MD&B 2027	HMD	Patients were able to explore five different virtual environments developed by Virtual Reality Medical Center in San Diago (cliff, dream castle, enchanted forest, icy cool world, and drive, walk, bike)	30 minutes	n/a	59 of 67 patients (88%) reported decreased pain level experienced after VR. Pain intensity was decreased by 3.75 points (likert scale) on average, which corresponds to a change from "severe" to "moderate" or "moderate" to "light."	Pain immediately after VR.	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Walker, 2014 ³⁶	RCT	45	Range: 18-70	English speaking men referred for flexible cystoscopy.	MPRP	Acute	PEDro 7/10	Patients used a VR helmet and track ball hand controller as they underwent cystoscopy.	Patients engaged "SnowWorld," where they were tasked to shoot snowballs at the penguins, robots, and igloos.	During cyroscopy (exact length of time unspecified).	Routine cystoscopy, including intraurethral amni- nistration of 2% lidocaine jelly, and the ability to watch their cystoscopy on the monitor; and routine inter- time action with phy- sican without VR.	None of the measures of pain or anxiety showed improvement in the VR distraction group undergoing flex- ible cystoscopy (20% power). Additionally, there were no differences in VR post-prece- dural pain from patients who scored high pre- procedural pain scores (80-100mm). Patients reported not feeling immersed in VR en- rionment.	Average pain dur- ing pro- cedure	una- ble to cal- cula- te	unable to cal- culate	unable to cal- culate	
	—	—	—	—	—	—	—	—	—	—	—	—	Worst pain dur- ing pro- cedure	una- ble to cal- cula- te	unable to cal- culate	unable to cal- culate	unable to cal- culate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Jones, 2016 ⁴⁴	Quasi-experimental study (no comparison group)	30	Range: 18+	Adults with various chronic pain conditions lasting 1-43 years.	MSKPNP	Chronic	MD&B 18/27	Oculus Rift DK2 HMD with headphones and a handheld mouse	Patients engaged in "COOL!" wherein they are taken through a virtual landscape and can interact with aspects of the landscape by clicking on a mouse to toss orbs or toss fish. When hit, flames will make sounds and change colors. When hit, otters will move about in a playful way and change colors.	5 minutes	n/a	The average pre-session pain rating for the 30 participants was 5.7. The average pain rating during the VR session was 2.6. The average change in pain rating between pre-session ratings and during-session ratings was -3.1 and resulted in a 60% reduction in pain. A paired t-test also found this change to be significant (p<0.001).	Pain during VR.	unable to calculate	unable to calculate	unable to calculate	

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
	—	—	—	—	—	—	—	—	—	—	—	The average pre-session pain rating for the 30 participants was 5.7. The average post-session pain rating was 4.1. The average change in pain rating between pre-session ratings and post-session ratings was -1.6 and resulted in a 33% reduction in pain. A paired t-test of pre-post session ratings found this change to be significant ($p<0.001$).	Pain immediately after VR.	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Harvie, 2015 ⁴²	RCT, within subjects cross over design	24	AVG 45	Average 11 years of chronic neck pain and pain with neck rotation, mildly to moderately disabled according to baseline NDI scores.	MSKP	Chronic	PEDro 8/10	Oculus Rift HMD with headphones Real-world movement was tracked and the fed into the virtual environment in an understated or overstated form. Participants sat in supportive chairs that prevented trunk movement.	Patients engaged in six scenes including four outdoor scenes (a park, a mountain, a countryside, and church grounds) and two indoor scenes (a dining room and a living room).	Participants rotated their head slowly to left and stop at pain, then right and stop at pain, for all three conditions.	20% less than (rotation gain = 0.8) actual physical rotation experienced during VR or 20% greater than (rotation gain = 1.2) actual physical rotation.	Pain-free ROM increased when visual feedback understated true rotation and decreased when visual feedback overstated true rotation (both significant results). No difference in pain intensity between all three conditions with VR (p = 0.6). No comparison made to non-VR condition.	Pain during head rotation task	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Powell, 2014 ⁴⁶	RCT, within subjects cross over design	19	AVG 54.8, range: 24-80	19 with musculoskeletal pain in upper or lower limb that compromised walking were recruited from the Jewish Rehabilitation Hospital (Laval, Quebec) and the Constance Lethbridge Rehabilitation Centre (Montreal, Quebec). 19 healthy volunteers were recruited from the staff and student body of the Jewish Rehabilitation Hospital, the Constance Lethbridge Rehabilitation Centre, and McGill University (Montreal, Quebec).	MSKP	Chronic	PEDro 5/10	Patients walked on a self-paced motorized treadmill. The treadmill responded dynamically to the speed of the user in real time. A 3D virtual walkway was created using Softimage XSI software and displayed on a large screen in front of the treadmill. Patients wore Logitech ClearChat wireless stereo headphones.	A 3D virtual walkway projected 2 parallel rows of vertical columns on either side of the walkway was displayed on the screen. The room was darkened, with the main light source being the display itself. Scene progression and audio feedback of footsteps on a hard surface were synchronized with treadmill speed.	12 x 2 minutes on treadmill	No audio; audio 75%, 100% or 125% of cadence. No comparison made to non-VR condition.	No significant changes in pain intensity ($p=0.65$) between beginning and end of experiment for the pain group. No comparison made to non-VR condition.	Pain immediately after VR.	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Sarig Bahat, 2015 ⁴⁷	RCT	32	VR group: 40.6 (SD 14.2), non-VR group: 41.1 (SD 12.6)	Participants were recruited in Brisbane, Australia. Participants were 18 y/o or greater, had neck pain > 3 months, and had an NDI score > 10%.	MSKP	Chronic	PEDro 7/10	HMD with 3D motion tracker built in (Wrap™ 1200VR by Vuzix)	Patients controlled a virtual pilot flying a red airplane via the patient's head motions and interacted with targets appearing from four directions (flexion, extension, right rotation, left rotation).	4-6 sessions for 30 minutes each over a 5 week period.	Kinematic training (KT) without VR. Both groups received KT.	NDI scores improved in both groups, but KT + VR group maintained this improvement at 3-months follow up. Only KT + VR group improved significantly in VAS at post-intervention.	Average Pain immediately after treatment	0.65	0.25	1.05	<0.05
	—	—	—	—	—	—	—	—	—	—	—	—	Average pain (in past week) - 3 months after intervention	0.51	0.14	0.88	NS

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Ichinose, 2017 ⁵⁰	Quasi-experimental study (no comparison group)	9	Range: 43-64	Brachial plexus avulsion injury and arm amputation, all patients perceived a phantom upper limb and pathological pain within it. (same sample as Osumi, 2016 plus one additional patient with arm amputation)	NP	Chronic	MD&B 17/27	Oculus Rift DK2 HMD. Spatial locations and movements of the intact arm and fingers were detected by two kinds of infrared video cameras (Kinect for Windows v2 and Leap Motion).	The virtual reality system would show bilateral limbs by mirroring the intact limb. Patients could have their virtual phantom limb reach for objects by moving their intact arm. When participants "touched" the object with the virtual phantom limb, an auditory ("collision" sound) and tactile (mechanical vibration) stimulus was provided. The vibrator was attached to the patient's cheek on their affected side or to their intact hand, or no vibrator was used.	15 minutes	n/a	Pain was significantly reduced in the Cheek (P = .004) and Intact Hand (P = .016) conditions. Statistical analyses showed that the pain reduction rates differed between conditions (Friedman test, $\chi^2 = 14.8$, P = .0006). The Cheek Condition was significantly higher than in the Intact Hand (P = .018) and No Stimulus (P = .0006) conditions. They calculated an "r values," which indicate the treatment effect sizes, were 0.68 (Cheek Condition vs No Stimulus Condition), 0.68 (Cheek Condition vs Intact Hand Condition), and 0.26 (Intact Hand Condition vs No Stimulus Condition).	Pain immediately after each VR condition.	unable to calculate	unable to calculate	unable to calculate	

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Osumi, 2017 ¹⁸	Quasi-experimental study (no comparison group)	8	Range: 43-64	Brachial plexus avulsion injury, all patients perceived a phantom upper limb and pathological pain within it.	NP	Chronic	MD&B 17/27	Oculus Rift DK2 HMD. Spatial locations and movements of the intact arm and fingers were detected by two kinds of infrared video cameras (Kinect for Windows v2 and Leap Motion).	The virtual reality system would show bilateral limbs by mirroring the intact limb. Patients could have their virtual phantom limb reach for objects by moving their intact arm. When participants "touched" the object with the virtual phantom limb, the object disappeared with a collision sound.	10 minutes	n/a	Short-term VR rehabilitation successfully and promptly alleviated phantom limb pain and simultaneously restored voluntary movement representations of a phantom limb. SF-MPQ averaged across all participants significantly decreased (pre 8.3 ± 7.6, post 2.5 ± 3.2, p=0.015). NRS pain scores decreased significantly (pre 5.2 ± 2.4, post 3.0 ± 2.1, p=0.015).	Pain immediately after VR.	0.98	0.29	1.66	0.02

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Wiederhold, 2014 ⁴³	RCT	31	Range: 18-65	Current non-cancer pain for at least 3 months and a daily average pain intensity score of 4.	NP	Chronic	PEDro 5/10	Delivery of the VR simulation was done with either a traditional HMD or a mobile phone display. A standard flat-panel display was used as a baseline. Half of the chronic pain participants received HMD exposure while the other half was exposed to a mobile device.	During exposure to the simulation patients were instructed to interact with the simulation graphics and explore the virtual worlds. No specifics were given on the exact virtual environment used.	20 minutes	Control group was described as a "pain focus" condition without VR.	All scales showed a subjective decrease in pain experienced while using both the mobile device and the HMD compared to the "pain focus" control condition. While the HMD was more effective in reducing subjective pain ratings, mobile devices also were able to achieve pain reduction effectively. The Visual Analog Pain Scale decreased when subjects were using the mobile phone pain distraction (mean score decreased by 0.58, p < 0.02), while it decreased by an additional 0.445 (p < 0.04) when the HMD was used.	Pain intensity during treatment	unable to calculate	unable to calculate	unable to calculate	

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Roosink, 2016 ⁴⁹	Quasi-experimental study (no comparison group)	9	Range: 25-72	A convenience sample was recruited among the outpatients of the Institut de réadaptation en déficience physique de Québec (IRDPQ). Individuals had sustained a traumatic SCI at least 3 months prior to participation, and had a lesion at the level of C4 or lower.	NP	Chronic	MD&B 17/27	The set-up consisted of an inertial movement sensor, a virtual reality system, two projectors (allowing for 3D vision), and a large silver-coated projection screen.	Patients engaged in a virtual forest path leading to the door of a small cabin (fixed distance) with a horizontal progress bar in the lower left corner of the screen (to provide feedback on the distance covered). Virtual walking consisted of 4 different conditions: forward with avatar, forward with static scene, backward with avatar, backward with static scene. The progress bar was updated in real-time based on right upper arm swing.	2 sessions 1.5 hours each, performed at least 1 hour apart.	n/a	Neuropathic pain intensity did not significantly change during the experiment. Reduced -2 on 0-100 scale and 95% confidence interval included 0 (-6 to 2).	Pain immediately after treatment	unable to calculate	unable to calculate	unable to calculate	unable to calculate

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Jin, 2016 ⁴⁵	RCT, within subjects cross over design	20	Range: 30-75	> 18 y/o with diagnosis of chronic pain (diagnosed by pain specialist). Recruited from complex pain clinic in Vancouver. 20% male, 80% female.	UnP	Chronic	PEDro 5/10	Oculus Rift DK2 HMD with noise-cancelling headphones and a computer mouse.	Patients engaged in the virtual reality game "Cryoslide" where they spent 4 minutes sliding in an icy cave and 6 minutes sliding in an outdoor icy world while throwing snowballs at creatures in the icy environment using a mouse.	10 minutes	Self-mediated pain control for 10 minutes (engage in activities they normally use to distract from pain such as meditating, reading, etc).	Significant reduced pain intensity in VR group compared to control group during the interventions (36.7% reduction, p < 0.001).	Pain during treatment	unable to calculate	unable to calculate	unable to calculate	
	—	—	—	—	—	—	—	—	—	—	—	No difference in pain intensity in VR group compared to control group after the interventions (p=0.265).	Pain immediately after treatment (within 10 minutes).	unable to calculate	unable to calculate	unable to calculate	

(Continued)

Table 1 (Continued)

Reference	Study Design	Sample Size	Age of participants (years)	Study Population	Type of Pain Condition	Stage of Pain Condition	Quality Assessment Score	VR Equipment	VR Intervention/ Environment	VR Dosage	Comparison Intervention	Main Results	Outcome Measure	Effect Size	95% CI (lower bounds)	95% CI (upper bounds)	Statistical Significance
Wiederhold, 2014 ⁴³	Quasi-experimental study (no randomization reported), within subjects design	40	Range: 22-68	Average daily pain >4/10 for >3 months	UnP	Chronic	MD&B 18/27	HMD with 3D motion tracker built in (Wrap™ 1200VR by Vuzix)	Patients engaged in pleasant and relaxing scenes such as forests, beaches, and mountains, relaxing music, and soothing effects such as the branches swaying and tall grass moving.	15 minutes	No distraction, described as a "pain focus" condition without VR.	All participants, reported a decrease in pain while in the VE ($p<0.05$).	Pain intensity after treatment (timing unreported)	unable to calculate	unable to calculate	unable to calculate	unable to calculate

Note: * $P<0.05$.

Abbreviations: VR, virtual reality; TBSA, total body surface area; RCT, randomized controlled trial; AVG, average; NDI, neck disability index; BP, burn pain; MPRP, medical-procedure related pain; MSKP, musculoskeletal pain; NP, neuropathic pain; MSKP-NP, musculoskeletal and neuropathic pain; UnP, unspecified pain; MD&B, Modified Downs and Black; NRS, numeric rating scale; SF-MPQ, short-form McGill Pain Questionnaire; KT, kinematic training; IVR, immersive virtual reality; NS, nonsignificant; GRS, graphic rating scale.

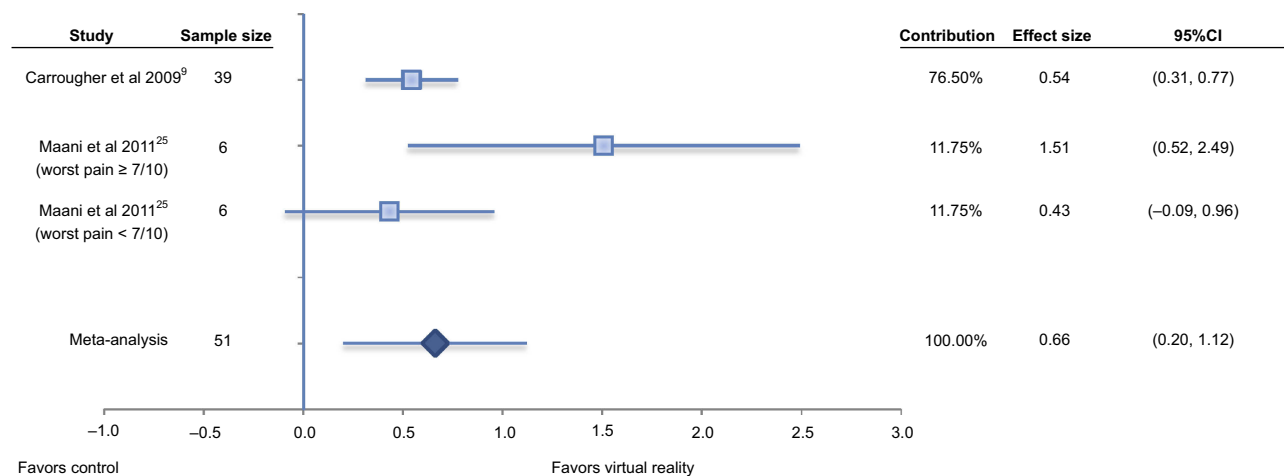


Figure 2 Effect size of virtual reality of reducing pain intensity during burn treatments (wound management or remobilization of limbs) compared to medication alone.

The third study found VR significantly reduced the worst pain experienced during labor contractions compared to no VR (ES = unable to calculate).³⁷ The fourth study found no significant change in either average or worst pain felt during a cystoscopy procedure with the addition of VR (ES = unable to calculate).³⁶ However, this study found that patients reported a sense of not feeling immersed in the VR environment. Meta-analysis was not performed due to the heterogeneity of these patient populations and medical procedures. All four of these studies used a HMD for VR application and had a dosage of 10 minutes,³⁷ 11 minutes,³⁵ 20 minutes³⁴ or the length of the procedure, which was not specified.³⁶ Quality assessment scores for these studies were between 6–7 and 10 on the PEDro scale and 22 and 27 on the MD&B scale indicating high quality and low risk of bias.

Impact of VR on pain experienced after procedure

Four studies evaluated the impact of VR on pain after various MPRP.^{35,38–40} The first of these studies reported a reduction in pain with the addition of VR during an episiotomy repair both immediately after (ES=1.37, 95%CI: 0.95, 1.79) and one hour after (ES=0.62, 95%CI: 0.34, 0.90) the procedure compared to standard episiotomy without VR.³⁵ The second study found significantly lower pain levels (ES=1.87, 95%CI: 1.49, 2.25) after a dressing change for hand injury when using VR compared to the dressing change with no VR.³⁸ The third study found no significant difference between VR after dressing change for various types of wounds (burns, ulcers, necrotizing fasciitis), compared to dressing change with no VR.⁴⁰ Interestingly, we calculated a significant ES=0.41 (95%CI: 0.09, 0.74), as this statistic is less impacted by small sample size compared to *P*-values.⁴¹ Meta-analysis

was not performed due to the heterogeneity of these patient populations and medical procedures. All of these studies used an HMD for VR application and had dosage from 11 minutes to approximately 30 minutes. Quality assessment scores for these studies were between 6–7 out of 10 on the PEDro scale indicating high quality and low risk of bias. The fourth study to look at pain after a medical procedure was a quasi-experimental study in patients within 24 hours of a cardiac procedure and found that the majority of patients (88%) experienced less pain immediately after a 30-minute application of VR with an HMD.³⁹ The quality assessment score for this study was 20/27 on the MD&B scale indicating low risk of bias.

Chronic pain conditions

Impact of VR on pain experienced during VR exposure

One study examined the effect of various forms of VR manipulation on chronic neck pain (MSKP) during the VR exposure.⁴² Specifically, they compared VR with active neck rotation motions to the onset of pain such that the VR environment either matched, underrepresented (by 20%), or overrepresented (by 20%) the actual neck motion. They did not compare to a nonR condition. They found that pain free range of motion increased with the VR experience that underrepresented the actual neck motion but that there was no difference in neck pain intensity experienced between the three conditions (ES = unable to calculate). This study used an HMD with a single rotation to each side for the three conditions for a total of six motions with VR. The PEDro score for this study was 8/10 indicating high quality and low risk of bias.

One study examined chronic NP with 20 minutes exposure to VR using an HMD or a mobile phone display compared to a control condition described as “pain focus” without VR.⁴³ This study found a significant reduction in pain during the VR exposure for both the HMD and mobile device compared to the control condition (ES = unable to calculate). The HMD led to a greater reduction in pain than the mobile device. The PEDro score for this study was 5/10 indicating a fair quality rating.

Another study examined a 5-minute exposure to VR using an HMD in patients with chronic MSKP-NP conditions but did not compare to a comparison group (quasi-experimental study).⁴⁴ This study found a significant decrease in pain within this group during the VR exposure by an average of 60% ($P<0.001$). The MD&B score was 18/27 indicating low risk of bias.

Finally, another study examined patients with UnP conditions.⁴⁵ In this study, they compared a 10-minute exposure of VR using an HMD to 10 minutes of self-mediated pain control where patients were instructed to engage in activities that they would normally use to distract them from their pain. They found that VR reduced pain intensity during the VR exposure by 36.7% ($P<0.001$) (ES = unable to calculate). The PEDro score for this study was 5/10 indicating a fair quality rating.

Impact of VR on pain experienced after VR exposure

Two studies examined pain intensity after VR exposure in patients with MSKP.^{46,47} The first of these studies projected a VR environment onto a screen in front of a treadmill where the participant walked at a self-selected speed for 12 bouts of 2 minutes with the audio feedback being either absent or present with a 75%, 100% or 125% match of their walking cadence.⁴⁶ They did not compare to a nonVR condition. They found that the VR experience with audio feedback provided at 125% of their baseline cadence led to an increase walking speed but that there was no significant change in pain intensity immediately after the VR exposures (ES = unable to calculate). The PEDro score for this study was 5/10 indicating a fair quality rating. The second study compared VR using HMD for 4–6 sessions of 30 minutes (over 5 weeks) in conjunction with kinematic training (KT) of active head and neck movements compared to the KT without VR.⁴⁷ They found that the VR group had a significant reduction in pain immediately after the VR exposure ($P<0.05$) (ES=0.65, 95%CI: 0.25, 1.05) but that this was not maintained at a 3-month follow-up (ES=0.51, 95%CI: 0.14, 0.88). They also noted significant

reductions in the Neck Disability Index (NDI) both immediately after VR exposure and at the 3-month follow-up ($P<0.01$ for both). The PEDro score for this study was 7/10 indicating high quality and low risk of bias.

Three studies examined pain intensity after VR exposure in NP conditions using a quasi-experimental design.^{48–50} The first of these studies evaluated a 10-minute VR exposure using HMD in patients with brachial plexus avulsion injuries that resulted in phantom limb pain.⁴⁸ A subsequent paper by the same authors reported on this same sample with the addition of one additional patient with an arm amputation under slightly different VR testing conditions.⁵⁰ Osumi et al⁴⁸ found that pain was significantly reduced immediately after 10 minutes of VR exposure ($P=0.015$). Similarly, Ichinose et al⁵⁰ found that pain was significantly reduced immediately after 15 minutes of VR exposure that included tactile stimulation on the patient’s cheek that coincided with a task within the VR environment ($P=0.004$). The MD&B score was 17/27 for both studies indicating low risk of bias. The third study evaluated two sessions of 1.5 hours (≥ 1 hour apart) of a VR environment projected onto two screens in front of patients with a spinal cord injury (C4 or lower) at least 3 months prior.⁴⁹ This study found no significant difference in pain reported immediately after the VR exposure. The MD&B score was 17/27 indicating low risk of bias.

Another study examined a 5-minute exposure to VR using an HMD in patients with chronic MSKP-NP conditions but did not compare to a comparison group (quasi-experimental study).⁴⁴ This study found a significant decrease in pain within this group immediately after the VR exposure by an average of 33% ($P<0.001$). The MD&B score was 18/27 indicating low risk of bias.

Two studies examined the impact of VR on pain intensity after VR exposure in patients with chronic UnP conditions.^{43,45} The first study compared a 10-minute exposure of VR using an HMD to a self-mediated pain control where patients were instructed to engage in activities that they would normally use to distract from their pain for 10 minutes without VR.⁴⁵ They found no difference in pain intensity between groups within 10 minutes of completion of the VR exposure ($P=0.265$) (ES = unable to calculate). The PEDro score for this study was 5/10 indicating a fair quality rating. The other study evaluated 15 minutes exposure to VR using an HMD compared to a control condition described as “pain focus” without VR (within subjects design, no randomization reported).⁴³ They found a significant reduction in pain intensity after VR exposure (time frame not reported) ($P<0.05$) (ES =

unable to calculate). The MD&B score was 18/27 indicating low risk of bias.

Discussion

The majority of the acute pain studies showed that VR is an effective tool in reducing acute pain experienced during various medical procedures (labor contractions, episiotomy repair, periodontal procedures) or burn-related pain (wound debridement or remobilization of range of motion for joints limited by burns). Only one study of acute pain found no difference in pain experienced during a cystoscopy procedure with the addition of VR, however, the authors reported the patients did not feel immersed in the VR environment.³⁶

In comparison, three of the four acute pain studies that looked at pain experienced after various medical procedures (episiotomy repair, dressing changes for hand injuries, and cardiac procedures) found a significant reduction in pain with the addition of VR. One study did not find a significant difference in pain levels after wound care with VR compared to no VR. However, we calculated a moderate effect size in favor of VR, indicating that this study may have been underpowered to detect this level of effect. All of these studies were rated as high quality and low risk of bias. These findings suggest that there is significant evidence to support the use of VR to reduce the severity of pain experienced both during and after many medical procedures. The magnitude of pain reduction varies depending on patient population, pain condition, and timing of measurement.

The findings for the chronic pain studies are less consistent. The two experimental trials in patients with chronic NP or UnP that compared to a nonVR comparison group (a “pain focus” condition or a self-mediated pain control session, respectively) found that pain intensity was significantly reduced during the VR exposure.^{43,45} The quality assessment for these two studies was only fair indicating some potential risk of bias in the study design. Further support for reductions in pain intensity during VR exposure in patients with chronic pain comes from a study of patients with MSKP-NP.⁴⁴ In contrast, another study looking at various forms of VR experiences in MSKP found no difference in neck pain during three versions of VR exposure.⁴² Although the two aforementioned studies had low risk of bias, they did not compare to a nonVR condition which limits the ability to evaluate the impact of VR compared to other forms of intervention for chronic pain.

Two experimental trials in patients with either chronic MSKP⁴⁷ or chronic UnP⁴³ found reduced pain after VR exposure compared to nonVR comparisons (kinematic training of active head and neck movements or a self-mediated pain control session, respectively). The former study found that reductions in pain level were not maintained at 3 months after VR exposure. In contrast, one study found no significant difference in pain intensity within 10 minutes after VR exposure compared to a “pain focus” condition in patients with UnP.⁴⁵ A similar pattern was noted from quasi-experimental studies, such that three studies found a reduction of pain after VR exposure in patients with NP or MSK-NP and one study found no difference in patients with NP. Lastly, another study looking at various forms of VR experiences in MSKP found no difference in pain experienced immediately after VR exposure, however, they did not compare to a nonVR condition, which limits the ability to evaluate the impact of VR compared to other forms of intervention for chronic pain.⁴⁶

All chronic pain studies had fair to high quality assessment ratings. The findings of the chronic pain studies suggest that there is potential for VR to reduce pain during VR exposure in patients with chronic pain. However, there are inconsistent findings for the use of VR to reduce pain after VR exposure in patients with chronic pain. Therefore, it may be speculated that although VR may effectively reduce chronic pain intensity during and possibly immediately after the VR exposure, there is unlikely to be a lasting analgesic effect in patients with chronic pain conditions with the treatment protocols tested to date.

One possibility for the variance found in our chronic pain results is the dosage of VR intervention. For example, when patients underwent a single 10-minute bout of VR, the VR group had no significant reduction in their pain.⁴⁵ However, when patients underwent 4–6×30-minute sessions over a 5-week period, the VR group had a significant reduction in pain immediately after VR exposure.⁴⁷ This may indicate the need for a larger dose of VR therapy to impact pain for people with chronic pain conditions, but requires further research.

A second possibility may be related to the type of VR equipment used. The majority of studies used a form of glasses/goggles or other forms of HMD, while two studies used projectors. The two studies that used projectors did not support the use of VR to alter chronic pain. Therefore, it may be possible that the use of an HMD allows for a

greater level of immersion and thus more distraction from the user's pain than a projector.

Our findings are consistent with previous systematic reviews, which have found VR to be effective in reducing acute pain.^{18,19,21,51} However, our systematic review includes more research on pain intensity and makes a direct comparison between changes in acute and chronic pain. One systematic review concluded that VR distraction was ineffective in reducing chronic pain, however, this conclusion was based on only one study.¹⁹ We were able to compare 10 chronic pain articles and found new evidence to suggest that VR intervention has the potential to significantly reduce chronic pain during VR intervention. The most recent systematic review¹⁹ did not provide quality assessment scores for included articles, which limits the reader's ability to assess the validity of their findings.

This systematic review was focused on the effects of VR intervention on reducing pain intensity, but other studies have looked at the effects of VR on functional outcomes. For example, Sarig Bahat et al⁴⁷ measured the impact of chronic neck pain on functional performance using the NDI and found that the VR group had positive changes in NDI scores for 3 months postintervention compared to the control group. It will be important to further examine the effect of VR on function in future research.

One strength of our review was the broad search strategy including three databases and searching other systematic reviews on the same topic for any studies that met our criteria. Previous studies^{19,27} have focused on experimental pain or have focused solely on pediatric populations. We excluded experimental pain to promote clinical relevance of our findings and excluded pediatric populations to improve our study's applicability to adults.

There was significant heterogeneity in study population and pain conditions, which poses a limitation for this systematic review. Additionally, our review is limited by the variety of the VR equipment used and the dosage of VR intervention in the included studies. One other limitation is that we included articles written in English only.

As VR continues to emerge as an adjunctive pain reduction treatment, it is important to consider the cost, feasibility and accessibility of VR equipment. The VR equipment in this review ranged from a high cost multiple-projector set-up to a low-cost commercially available system and the VR market is predicted to grow 29.2% by 2020.⁵² With this evolving market, it is an ethical consideration to ensure that access to VR treatment does not become cost-prohibitive and that application of this

intervention is supported by the literature. Future studies should continue to examine the cost-effectiveness of VR as a pain management tool, and if it is feasible to use low-cost VR systems to reduce the financial burden on the health-care system and improve access.

In addition, this review highlights the need for high quality randomized clinical trials for VR in patients with chronic pain conditions. It is recommended that studies evaluate appropriate dosage of VR exposure compared to other pain analgesia therapies (ie pain medications, visual illusions, mirror box therapy, etc.) for both short and long-term outcomes that are meaningful to the patient (including pain reduction, function and quality of life). Moreover, future studies should examine the effects of VR on pain control in different clinical settings (including intensive care, acute care and outpatient settings).

One additional recommendation comes from the considerable variability in the definition of immersive VR across studies. A possible solution is to provide a standardized measurement of how immersed participants feel when undergoing VR intervention. Some of the studies included in our review used subjective measures of immersiveness to quantify their experience of immersion. For example, participants rated their sense of presence in the virtual environment on a VAS scale from, "I did not feel like I went inside at all," to, "I went completely into the computer-generated world."⁹ However, we believe a more comprehensive tool is warranted due to the complexity of immersion. One of our included studies¹² used the Presence Questionnaire (PQ) developed by Witmer and Singer.⁵¹ The PQ is a 24-question, valid and reliable measure of presence in a virtual world.⁵³ We recommend researchers consider the PQ for its ability to capture multiple dimensions of the patient's experience.

Finally, it may also be imperative to determine if there is a clinical difference between "immersion" and "presence." "Presence" may be defined as the subjective illusion in the participant's mind that he or she has gone inside the virtual world.^{19,51} For example, the person can receive sensory feedback from the virtual world, but may not be "present" if they are not cognitively engaged in it. Previous studies have attempted to explain VR's effect on pain, noting that decreased activation of pain processing areas in the brain may be responsible for subjective pain reduction with VR intervention.^{7,51,54} A shift in cognitive demand can lead to distraction from pain, and being "present" in the virtual environment can theoretically stimulate this shift. Even if someone is "immersed" in a virtual world, they may not be

“present” in it. We recommend further research to determine if “immersion” and “presence” are conceptually different, and if so what impact they have on VR in reducing pain.

Conclusion

This study provides an updated systematic literature review showing the effects of VR on both acute and chronic pain. We conclude that VR is an effective tool in reducing acute pain both during and after a VR intervention, and that VR is especially effective in mitigating MPRP. The evidence for chronic pain relief with VR suggests that chronic pain may be reduced while the patient is immersed in the VR environment, but with minimal long-term carryover effect beyond the immediate postVR exposure time frame. Further research is needed to assess the extent to which one needs to be immersed and present in a virtual environment in order to reduce pain, and the dosage necessary to maintain pain reductions in chronic pain over time. Clinicians should consider immersive VR therapies as an adjunct to²³ standard care to help reduce acute pain and potentially for chronic pain conditions.

Acknowledgments

The authors are grateful to Dr Jason Hardage for his assistance with our review process and to Meira Prescher, Ping Zhang, Eric Kwan, and AJ Pluss for their thorough peer review. No sources of funding were used to create this systematic review.

Disclosure

The authors report no conflicts of interest in this work.

References

1. Rasu RS, Vouthy K, Crowl AN, et al. Cost of pain medication to treat adult patients with nonmalignant chronic pain in the United States. *J Manag Care Spec Pharm*. 2014;20(9):921–928. doi:10.18553/jmcp.2014.20.9.921
2. Teater D The psychological and physical side effects of pain medications. 2014. Available from: <http://safety.nsc.org/sideeffects>. Accessed September 25, 2018.
3. Sutherland S Meeting report: virtual reality pain 2016 | pain research forum. 2016. Available from: <https://www.painresearchforum.org/news/74439-meeting-report-virtual-reality-pain-2016>. Accessed September 25, 2018.
4. Greenleaf W How virtual and augmented reality will change the way we shop | techno FAQ. Available from: https://www.slideshare.net/waltergreenleaf/how-virtual-and-augmented-reality-will-transform-healthcare-august-2017-presentation?qid=16451079-9e92-44a7-9baf-1e1d237c8575&v=&b=&from_search=1. Accessed September 25, 2018.
5. Hoffman HG, Doctor JN, Patterson DR, Carrougner GJ, Furness TA. Virtual reality as an adjunctive pain control during burn wound care in adolescent patients. *Pain*. 2000;85(1–2):305–309.
6. Hoffman HG, Patterson DR, Carrougner GJ, Sharar SR. Effectiveness of virtual reality-based pain control with multiple treatments. *Clin J Pain*. 2001;17(3):229–235. doi:10.18553/jmcp.2014.20.9.921
7. Hoffman HG, Seibel EJ, Richards TL, Furness TA, Patterson DR, Sharar SR. Virtual reality helmet display quality influences the magnitude of virtual reality analgesia. *J Pain*. 2006;7(11):843–850. doi:10.1016/j.jpain.2006.04.006
8. Hoffman HG, Richards TL, Van Oostrom T, et al. The analgesic effects of opioids and immersive virtual reality distraction: evidence from subjective and functional brain imaging assessments. *Anesth Analg*. 2007;105(6):1776–1783. doi:10.1213/01.ane.0000270205.45146.db
9. Carrougner GJ, Hoffman HG, Nakamura D, et al. The effect of virtual reality on pain and range of motion in adults with burn injuries. *J Burn Care Res*. 2009;30(5):785–791. doi:10.1097/BCR.0b013e3181b485d3
10. Hoffman HG, Chambers GT, Meyer WJ, et al. Virtual reality as an adjunctive non-pharmacologic analgesic for acute burn pain during medical procedures. *Ann Behav Med*. 2011;41(2):183–191. doi:10.1007/s12160-010-9248-7
11. Schmitt YS, Hoffman HG, Blough DK, et al. A randomized, controlled trial of immersive virtual reality analgesia, during physical therapy for pediatric burns. *Burns*. 2011;37(1):61–68. doi:10.1016/j.burns.2010.07.007
12. Schneider SM, Hood LE. Virtual reality: a distraction intervention for chemotherapy. *Oncol Nurs Forum*. 2007;34(1):39–46. doi:10.1188/07.ONF.39-46
13. Schneider SM, Ellis M, Coombs WT, Shonkwiler EL, Folsom LC. Virtual reality intervention for older women with breast cancer. *Cyberpsychol Behav*. 2003;6(3):301–307. doi:10.1089/109493103322011605
14. Windich-Biermeier A, Sjoberg I, Dale JC, Eshelman D, Guzzetta CE. Effects of distraction on pain, fear, and distress during venous port access and venipuncture in children and adolescents with cancer. *J Pediatr Oncol Nurs*. 2007;24(1):8–19. doi:10.1177/1043454206296018
15. Sharar SR, Miller W, Teeley A, et al. Applications of virtual reality for pain management in burn-injured patients. *Expert Rev Neurother*. 2008;8(11):1667–1674. doi:10.1586/14737175.8.11.1667
16. Miller K, Rodger S, Bucolo S, Greer R, Kimble RM. Multi-modal distraction. Using technology to combat pain in young children with burn injuries. *Burns*. 2010;36(5):647–658. doi:10.1016/j.burns.2009.06.199
17. Garrett B, Taverner T, Masinde W, Gromala D, Shaw C, Negraeff M. A rapid evidence assessment of immersive virtual reality as an adjunct therapy in acute pain management in clinical practice. *Clin J Pain*. 2014;30(12):1089–1098. doi:10.1097/AJP.0000000000000064
18. Shahrbanian S, Ma X, Korner-Bitensky N, Simmonds MJ. Scientific evidence for the effectiveness of virtual reality for pain reduction in adults with acute or chronic pain. *Stud Health Technol Inform*. 2009;144:40–43.
19. Malloy KM, Milling LS. The effectiveness of virtual reality distraction for pain reduction: a systematic review. *Clin Psychol Rev*. 2010;30(8):1011–1018. doi:10.1016/j.cpr.2010.07.001
20. Macedonio MF, Parsons TD, DiGiuseppe RA, Weiderhold BK, Rizzo AA. Immersiveness and physiological arousal within panoramic video-based virtual reality. *Cyberpsychol Behav*. 2007;10(4):508–515. doi:10.1089/cpb.2007.9997
21. Chan E, Foster S, Sambell R, Leong P. Clinical efficacy of virtual reality for acute procedural pain management: a systematic review and meta-analysis. *PLoS One*. 2018;13(7):e0200987. doi:10.1371/journal.pone.0200987
22. Gupta A, Scott K, Dukewich M. Innovative technology using virtual reality in the treatment of pain: does it reduce pain via distraction, or is there more to it? *Pain Med*. 2018;19(1):151–159. doi:10.1093/pm/pnx109

23. Treede R-D, Rief W, Barke A, et al. A classification of chronic pain for ICD-11. *Pain*. 2015;156(6):1. doi:10.1097/j.pain.000000000000160
24. Boesch E, Bellan V, Moseley GL, Stanton TR. The effect of bodily illusions on clinical pain. *Pain*. 2016;157(3):516–529. doi:10.1097/j.pain.0000000000000423
25. Maani CV, Hoffman HG, Morrow M, et al. Virtual reality pain control during burn wound debridement of combat-related burn injuries using robot-like arm mounted VR goggles. *J Trauma*. 2011;71(1 Suppl):S125–S130. doi:10.1097/TA.0b013e31822192e2
26. Hoffman HG, Patterson DR, Carrouger GJ. Use of virtual reality for adjunctive treatment of adult burn pain during physical therapy: a controlled study. *Clin J Pain*. 2000;16(3):244–250.
27. Wender R, Hoffman HG, Hunner HH, Seibel EJ, Patterson DR, Sharar SR. Interactivity influences the magnitude of virtual reality analgesia. *J Cyber Ther Rehabil*. 2009;2(1):27–33.
28. Neyeloff JL, Fuchs SC, Moreira LB. Meta-analyses and Forest plots using a microsoft excel spreadsheet: step-by-step guide focusing on descriptive data analysis. *BMC Res Notes*. 2012;5(1):52. doi:10.1186/1756-0500-5-52
29. de Morton NA. The PEDro scale is a valid measure of the methodological quality of clinical trials: a demographic study. *Aust J Physiother*. 2009;55(2):129–133.
30. Downs SH, Black N. The feasibility of creating a checklist for the assessment of the methodological quality both of randomised and non-randomised studies of health care interventions. *J Epidemiol Community Health*. 1998;52(6):377–384. doi:10.1136/jech.52.6.377
31. Teasell RW, Foley NC, Bhogal SK, Speechley MR. An evidence-based review of stroke rehabilitation. *Top Stroke Rehabil*. 2003;10(1):29–58. doi:10.1310/8YNA-1YHK-YMHB-XTE1
32. Abou El Fadl R, Blair M, Hassounah S. Integrating maternal and children's oral health promotion into nursing and midwifery practice—a systematic review. *PLoS One*. 2016;11(11):e0166760. doi:10.1371/journal.pone.0166760
33. Morris LD, Louw QA, Crous LC. Feasibility and potential effect of a low-cost virtual reality system on reducing pain and anxiety in adult burn injury patients during physiotherapy in a developing country. *Burns*. 2010;36(5):659–664. doi:10.1016/j.burns.2009.09.005
34. Furman E, Jasinevicius TR, Bissada NF, Victoroff KZ, Skillicorn R, Buchner M. Virtual reality distraction for pain control during periodontal scaling and root planing procedures. *J Am Dent Assoc*. 2009;140(12):1508–1516.
35. JahaniShoorab N, Ebrahimzadeh Zagami S, Nahvi A, et al. The effect of virtual reality on pain in primiparity women during episiotomy repair: a randomized clinical trial. *Iran J Med Sci*. 2015;40(3):219–224.
36. Walker MR, Kallungal GJS, Musser JE, Folen R, Stetz MC, Clark JY. Treatment efficacy of virtual reality distraction in the reduction of pain and anxiety during cystoscopy. *Mil Med*. 2014;179(8):891–896. doi:10.7205/MILMED-D-13-00343
37. Frey DP, Bauer ME, Bell CL, et al. Virtual reality analgesia in labor. *Anesth Analg*. 2018;1. doi:10.1213/ANE.0000000000003649
38. Guo C, Deng H, Yang J. Effect of virtual reality distraction on pain among patients with hand injury undergoing dressing change. *J Clin Nurs*. 2015;24(1–2):115–120. doi:10.1111/jocn.12626
39. Mosso-Vázquez JL, Gao K, Wiederhold BK, Wiederhold MD. Virtual reality for pain management in cardiac surgery. *Cyberpsychol Behav Soc Netw*. 2014;17(6):371–378. doi:10.1089/cyber.2014.0198
40. McSherry T, Atterbury M, Gartner S, Helmold E, Searles DM, Schulman C. Randomized, crossover study of immersive virtual reality to decrease opioid use during painful wound care procedures in adults. *J Burn Care Res*. 2018;39(2):278–285. doi:10.1097/BCR.0000000000000589
41. Sullivan GM, Feinn R. Using effect size-or why the p-value is not enough. *J Grad Med Educ*. 2012;4(3):279–282. doi:10.4300/JGME-D-12-00156.1
42. Harvie DS, Broecker M, Smith RT, Meulders A, Madden VJ, Moseley GL. Bogus visual feedback alters onset of movement-evoked pain in people with neck pain. *Psychol Sci*. 2015;26(4):385–392. doi:10.1177/0956797614563339
43. Wiederhold BK, Gao K, Sulea C, Wiederhold MD. Virtual reality as a distraction technique in chronic pain patients. *Cyberpsychol Behav Soc Netw*. 2014;17(6):346–352. doi:10.1089/cyber.2014.0207
44. Jones T, Moore T, Choo J. The impact of virtual reality on chronic pain. *PLoS One*. 2016;11(12):e0167523. doi:10.1371/journal.pone.0167523
45. Jin W, Choo A, Gromala D, Shaw C, Squire P. A virtual reality game for chronic pain management: a randomized, controlled clinical study. *Stud Health Technol Inform*. 2016;220:154–160.
46. Powell W, Simmonds MJ. Virtual reality and musculoskeletal pain: manipulating sensory cues to improve motor performance during walking. *Cyberpsychol Behav Soc Netw*. 2014;17(6):390–396. doi:10.1089/cyber.2014.0061
47. Sarig Bahat H, Takasaki H, Chen X, Bet-Or Y, Treleaven J. Cervical kinematic training with and without interactive VR training for chronic neck pain - a randomized clinical trial. *Man Ther*. 2015;20(1):68–78. doi:10.1016/j.math.2014.06.008
48. Osumi M, Ichinose A, Sumitani M, et al. Restoring movement representation and alleviating phantom limb pain through short-term neurorehabilitation with a virtual reality system. *Eur J Pain*. 2017;21(1):140–147. doi:10.1002/ejp.910
49. Roosink M, Robitaille N, Jackson PL, Bouyer LJ, Mercier C. Interactive virtual feedback improves gait motor imagery after spinal cord injury: an exploratory study. *Restor Neurol Neurosci*. 2016;34(2):227–235. doi:10.3233/RNN-150563
50. Ichinose A, Sano Y, Osumi M, Sumitani M, Kumagaya S-I, Kuniyoshi Y. Somatosensory feedback to the cheek during virtual visual feedback therapy enhances pain alleviation for phantom arms. *Neurorehabil Neural Repair*. 2017;31(8):717–725. doi:10.1177/1545968317718268
51. Witmer BG, Singer MJ. Measuring presence in virtual environments: a presence questionnaire. *Pres Teleoperators Virtual Environ*. 1998;7(3):225–240. doi:10.1162/105474698565686
52. Kern C Dropping virtual reality prices yields better opportunities for healthcare. Health IT Outcomes website. Available from: <https://www.healthitoutcomes.com/doc/dropping-virtual-reality-prices-yields-better-opportunities-for-healthcare-0001>. Published 2016. Accessed September 25, 2018.
53. Dascal J, Reid M, IsHak WW, et al. Virtual reality and medical inpatients: a systematic review of randomized, controlled trials. *Innov Clin Neurosci*. 2017;14(1–2):14–21.
54. Li A, Montañó Z, Chen VJ, Gold JI. Virtual reality and pain management: current trends and future directions. *Pain Manag*. 2011;1(2):147–157. doi:10.2217/pmt.10.15

Journal of Pain Research

Dovepress

Publish your work in this journal

The Journal of Pain Research is an international, peer reviewed, open access, online journal that welcomes laboratory and clinical findings in the fields of pain research and the prevention and management of pain. Original research, reviews, symposium reports, hypothesis formation and commentaries are all considered for publication. The manuscript

management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit <http://www.dovepress.com/testimonials.php> to read real quotes from published authors.

Submit your manuscript here: <https://www.dovepress.com/journal-of-pain-research-journal>